

Case/Sample Summary Report

Date:

July 19, 2011

From:

James A. Turner

Lead Analyst, Organic Branch

Subject: Status Report

OCI Case #: 2010-PHP-715-0519; CIE Serial #: 165500

FACTS #: 677114

To:

Special Agent

Philadelphia Resident Office, OCI/FDA

Through: R. Duane Satzger, Ph.D.

Director, Organic Branch

This report presents the results for analyses that have been completed to date. Additional work is being done and a final report will be issued when that is finished.

I. Description of Samples Received for Analysis

The sample was received via UPS on 2/18/2011. The sample consisted of six items which were each described, in part, on CIE 165500. Additional description of the sample is provided in Table 1, below.

II. Analytical Tests Performed on the Samples

Item 3, Item 4, Item 5 and Item 6 were analyzed by one or more of the following techniques: Enzyme-Linked Immunosorbent Assay (ELISA), High Performance Liquid Chromatography / Mass Spectrometry (HPLC/MS), and HPLC/UV (High Performance Liquid Chromatography / Ultraviolet Detection) for purposes of identifying and quantifying drugs that were present. Additional information is provided in Table 1, below.

III. Results of the Analyses

The results of analyses are contained in Table 1 which is presented on pages 2 - 4.

Drugs were identified using HPLC/MS based upon a comparison of the retention time and mass spectral data of a component of the sample with those of a standard which was analyzed under the same conditions. Drugs were assayed using HPLC/UV.

For Item 3, the sample was analyzed using ELISA. The results were consistent with the presence of Human Chorionic Gonadotropin (hCG). This drug was declared on the labeling associated with the Item. This is indicated in Table 1.

The numbers in parentheses in Table 1 represent the 95% confidence interval about the reported assay values.

There was no evidence for the presence of any drugs other than those listed Table 1.



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Results of Analysis CIE 165500 (FACTS 677114)						Table 1
Sample		Description of Sample Technique				
Item 1	"Combipack of Mif Mifepristone200 following code "Rs B)A00439 MFD.FE code "Rs.499.00 N MFD.MAR.10 EXF	uct with corresponde epristone and Misoping B; 4 Misopro .499.00 MASTER B B.10 EXP.JUL.11" MASTER B. No.A00 P.AUG.11". Each bots and one Mifepristo	Analysis ongoing	Analysis ongoing		
Item 2	labeled, in part, "A vial with an intact of containing a white	roduct with corresponders and the control of the co				
		Falcigo Vial	Sodium Bicarbonate Ampoule	Sodium Chloride Ampoule	Analysis ongoing	Analysis ongoing
	Batch No. :	AFI1074	2690011	7210849		
	Mfg. Date :	09/2009	08/2009	06/2009		
	Expiry Date :	02/2011	07/2011	05/2014	1	1

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esults of	Analysis CIE 16550	00 (FACTS 677114)				Table 1 (continued)
Sample		Technique	Drug Identified / Assay Value			
Item 3	Five boxes of product with corresponding labeling and codes each of which was labeled, in part, "Chorionic Gonadotrophin Injection I.P. (Human Chorionic Gonadotrophin Injection)5000 I.U.". Each box held one clear vial with an intact green flip cap labeled, in part, as the box and containing a white powder and one intact amber ampoule labeled, in part, "Sodium Chloride Injection I.P." and containing a clear colorless free-flowing liquid. Two of five of the boxes/vials of product were analyzed – more specifically, only the Human Chorionic Gonadotropin (hCG) labeled vials were analyzed. Sodium hCG Vial Ampoule Batch No.: A66110004 A6610009			ELISA	The data is consistent with the presence of Human Chorionic Gonadotropin (hCG) (Assay work ongoing)	
	Mfg. Date : Expiry Date :	03/10 02/13	03/10 03/15			
Item 4	which was labeled MFG.AUG. 2010 B	l, in part, "Sildigra…S EXP, JULY 2012…".	corresponding labeling and cou sildenafil Citrate 100 mgB.No. Each blister-pack held ten grey of five blister-packs were analyz	SGSG 1004 unmarked soft-	HPLC/MS And HPLC/UV	Sildenafil at 72 (± 4) mg/capsule (This is equivalent to 102 (±6) mg/capsule as Sildenafil Citrate)

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esults of	Table 1 (continued)		
Sample	Description of Sample	Technique	Drug Identified / Assay Value
Item 5	Five intact blister-packs of product with corresponding labeling and codes each of which was labeled, in part, "FILAGRA 100Sildenafil Citrate Tablets 100 mgB.NO.T89012 M.D.OCT.2010 E.D.SEP 2013". Each blister-pack held ten diamond (bi-pyramid) shaped tablets with the following debossing "DP" and an oval shape on one side and "FGR-100" on the other side. Five tablets from one of five blister-packs were analyzed.	HPLC/MS And HPLC/UV	Sildenafil at 100 (<u>+</u> 8) mg/tablet
Item 6	Five intact blister-packs of product with corresponding labeling and codes each of which was labeled, in part, "TAdALiSTATadalafil 20 mg TabletsB.NO.798015 M.D.OCT.2010 E.D.MAR.2013". Each blister-pack held ten unmarked yellow oval shaped tablets. Five tablets from one of five blister-packs were analyzed.	HPLC/MS And HPLC/UV	Tadalafil at 19 (<u>+</u> 2) mg/tablet

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IV. Sample Retention/Disposition/Feedback Information

This evidence will be retained by the Forensic Chemistry Center pending notification of disposition from your office. If you have any questions, concerns or need additional information, please do not hesitate to contact me at (513) 679-2700 Ext. 2240, or you can contact Kevin J. Mulligan, Ph.D. at (513) 679-2700 Ext. 2238.

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Section Author's Concurrence

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Kevin & Mulligar, Ph. D.

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